

In the Claims:

Please replace all prior versions and listings of claims with the amended claims as follows:

1. **(Currently amended)** A method of identifying potential ligands for protein macromolecular targets comprising:

- (1) providing a set of models, wherein each model comprises three-dimensional structural information for a ligand or a ligand:macromolecule protein complex, wherein each ligand comprises a plurality of atoms and a plurality of bonds, wherein each model is related to the other models of the set by a an amino acid homologous structural feature of 25% or greater for proteins or a structural homology between the ligands of at least 4-6 atoms in common; and wherein the structural information is derived from physical observation and/or computational inference;
- (2) mapping spatial relationships between the models such that the models are superimposed with respect to the homologous structural feature;
- (3) identifying one or more pairs of matching bonds between ligands of the set, wherein the matching bonds comprise a bond of a first ligand (B1) and a bond of a second ligand (B2) that are superimposed in step (2) such that
 - (i) an atom at each end of the bond (B1) is within 1.8 angstrom of an atom at each end of the bond (B2),
 - (ii) the bond (B1) and the corresponding bond (B2) are of the same bond order, and
 - (iii) the bond (B1) and the corresponding bond (B2) are related by an angle of 30° or less;
- (4) selecting a plurality of subsets of atoms and/or bonds from each ligand; wherein each subset comprises a bond and/or, an atom connected to the matching bond as identified in (3);

- (5) generating and displaying output ligands, each output ligand comprising atoms and/or bonds of a first subset as selected in (4) and atoms and/or bonds of a second subset as selected in (4), wherein the first subset and the second subset comprise atoms and/or bonds derived from opposite ends of a matching bond as selected in (4).
2. **(Original)** The method of claim 1, wherein the output ligands comprise all atoms represented in the ligands of step (1).
3. **(Currently amended)** The method of claim 1, wherein each model of the set comprises a ligand:~~macromolecule~~ protein complex.
4. **(Original)** The method of claim 1, wherein one or more models of the set consist of a ligand.
5. **(Canceled)**
6. **(Currently amended)** The method of claim 1 5, wherein the protein macromolecule is a protein kinase, a G-protein coupled receptor, an immunoglobulin superfamily protein, a protease, or a zinc-finger containing protein.
7. **(Currently amended)** The method of claim 3, wherein each model of the set comprises an identical protein macromolecule.
8. **(Original)** The method of claim 1, wherein the structural information is derived from a physical observation.
9. **(Original)** The method of claim 3, wherein the structural information comprises information derived by a computational inference.
10. **(Currently amended)** The method of claim 1, wherein at least one ligand of (1) is a small molecule.
11. **(Original)** The method of claim 1, wherein the ligands are less than 1000 atomic mass units (a.m.u.).
12. **(Original)** The method of claim 1, wherein the ligands are less than 600 a.m.u.
13. **(Canceled)**

14. **(Currently amended)** The method of claim 1-13, wherein the structural homology comprises homology between a framework of the ligands.
15. **(Currently amended)** The method of claim 1-13, wherein the structural homology comprises homology between a pharmacophore model of the ligands.
16. **(Currently amended)** The method of claim 1-5, wherein the ~~macromolecule is a protein, and wherein~~ the homologous feature comprises structural homology between the proteins of at least 40% amino acid homology.
17. **(Canceled)**
18. **(Canceled)**
19. **(Currently amended)** The method of claim 16-17, wherein the homology comprises a shared polypeptide fold.
20. **(Original)** The method of claim 1, wherein the set comprises at least three models.
- 21-23. **(Canceled)**
24. **(Original)** The method of claim 1, further comprising the steps of:
 - (6) comparing output ligands of step (5) to the ligands of step (1); and
 - (7) storing output ligands that are not identical to the ligands used in a previous iteration of steps (2)-(5) in a machine-readable medium.
25. **(Original)** The method of claim 24, further comprising generating one or more output models, wherein each output model comprises the stored ligand docked into a target macromolecule.
26. **(Original)** The method of claim 25, further comprising refining the output models.
27. **(Original)** The method of claim 26, wherein the refining comprises performing energy minimization computations.
28. **(Original)** The method of claim 27, further comprising evaluating the output models.

29. **(Original)** The method of claim 28, further comprising assigning a score to each output model based on the evaluating.
30. **(Previously presented)** The method of claim 29, further comprising obtaining a composition comprising a compound including a ligand from a subset of output models, wherein the subset comprises output models having a score in a preselected range.
31. **(Original)** The method of claim 30, further comprising evaluating the composition.
32. **(Original)** The method of claim 31, wherein the evaluating comprises determining the ability of the compound to bind a target macromolecule, or the ability of the compound to modulate activity of a target macromolecule.
33. **(Original)** The method of claim 24, wherein steps 2-7 are repeated, and wherein the models superimposed in step (2) comprise the stored output ligands of step (7).
34. **(Original)** The method of claim 33, wherein the repeating is automatic.
35. **(Previously presented)** The method of claim 34, wherein the repeating stops when each ligand of step (7) is identical to a ligand mapped in the previous step (2) of the repetition.
36. **(Original)** The method of claim 1, wherein the structural information comprises hydrogen atoms of the ligands and the bonds to hydrogen atoms.
37. **(Original)** The method of claim 1, wherein the structural information does not comprise hydrogen atoms of the ligands.
38. **(Previously presented)** The method of claim 1, wherein the ligands comprise a macrocyclic moiety, and wherein at least two matching bonds between the ligands are identified within the macrocycle of each ligand.
39. **(Currently amended)** A method of identifying potential ligands for a protein macromolecular targets comprising:
 - (1) selecting a set of models from a plurality of models, wherein the selecting comprises identifying models comprising a an amino acid homologous structural feature of 25% or greater for proteins or a structural homology between the ligands of at

least 4-6 atoms in common, wherein each model comprises three-dimensional structural information for a ligand:~~macromolecule~~ protein complex, wherein each ligand comprises a plurality of atoms and a plurality of bonds; and
wherein the structural information is derived from physical observation and/or computational inference;

- (2) providing the set of models;
- (3) mapping spatial relationships between the models such that the models are superimposed with respect to the homologous structural feature;
- (4) identifying one or more pairs of matching bonds between ligands of the set, wherein the matching bonds comprise a bond of a first ligand (B1) and a bond of a second ligand (B2) that are superimposed in step (2) such that
 - (i) an atom at each end of the bond (B1) is within 1.8 angstrom of an atom at each end of the bond (B2),
 - (ii) the bond (B1) and the corresponding bond (B2) are of the same bond order, and
 - (iii) the bond (B1) and the corresponding bond (B2) are related by an angle of 30° or less;
- (5) selecting a plurality of subsets of atoms and/or bonds from each ligand; wherein each subset comprises a bond and/or, an atom connected to a matching bond as identified in (4);
- (6) generating and displaying output ligands, each output ligand comprising atoms and/or bonds of a first subset as selected in (5) and atoms and/or bonds of a second subset as selected in (5), wherein the first subset and the second subset comprise atoms and/or bonds derived from opposite ends of a matching bond as selected in (5).

42. (Currently amended) An apparatus comprising:

(a) a memory that stores executable instructions; and

(b) a processor that executes the instructions to:

(1) provide a set of models, wherein each model comprises three-dimensional structural information for a ligand or a ligand:~~macromolecule protein~~ complex, wherein each ligand comprises a plurality of atoms and a plurality of bonds, and

wherein each model is related to the other models of the set by a ~~an amino acid homologous structural feature of 25% or greater for proteins or a structural homology between the ligands of at least 4-6 atoms in common;~~ and

wherein the structural information is derived from physical observation and/or computational inference;

(2) map spatial relationships between the models such that the models are superimposed with respect to the homologous structural feature;

(3) identify one or more pairs of matching bonds between ligands of the set, wherein the matching bonds comprise a bond of a first ligand (B1) and a bond of a second ligand (B2) that are superimposed in step (2) such that

(i) an atom at each end of the bond (B1) is within 1.8 angstrom of an atom at each end of the bond (B2),

(ii) the bond (B1) and the corresponding bond (B2) are of the same bond order, and

(iii) the bond (B1) and the corresponding bond (B2) are related by an angle of 30° or less;

(4) select a plurality of subsets of atoms and/or bonds from each ligand;

wherein each subset comprises a bond and/or, an atom connected to the matching bond as identified in (3);

(5) generate output ligands, each output ligand comprising atoms and/or bonds of a first subset as selected in (4) and atoms and/or bonds of a second subset as selected in (4), wherein the first subset and the second subset comprise atoms and/or bonds derived from opposite ends of a matching bond as selected in (4);

(6) compare output ligands to the ligands of step (1);

(7) store output ligands that are not identical to the ligands of step (1);

(8) repeat steps (2)-(7), wherein the models superimposed in step (2) comprise the stored output ligands of step (7);

wherein the repeating stops when each output ligand of step (7) is identical to a ligand provided in the previous step (2) of the repetition.

43. **(Currently amended)** An article comprising machine-readable media that stores executable instructions, the instructions causing a machine to:

(1) provide a set of models, wherein each model comprises three-dimensional structural information for a ligand or a ligand:macromolecule protein complex, wherein each ligand comprises a plurality of atoms and a plurality of bonds;

wherein each model is related to the other models of the set by a an amino acid homologous structural feature of 25% or greater for proteins or a structural homology between the ligands of at least 4-6 atoms in common; and

wherein the structural information is derived from physical observation and/or computational inference;

(2) map spatial relationships between the models such that the models are superimposed with respect to the homologous structural feature;

- (3) identify one or more pairs of matching bonds between ligands of the set, wherein the matching bonds comprise a bond of a first ligand (B1) and a bond of a second ligand (B2) that are superimposed in step (2) such that
- (i) an atom at each end of the bond (B1) is within 1.8 angstrom of an atom at each end of the bond (B2),
 - (ii) the bond (B1) and the corresponding bond (B2) are of the same bond order, and
 - (iii) the bond (B1) and the corresponding bond (B2) are related by an angle of 30° or less;
- (4) select a plurality of subsets of atoms and/or bonds from each ligand; wherein each subset comprises a bond and/or, an atom connected to the matching bond as identified in (3);
- (5) generate and display output ligands, each output ligand comprising atoms and/or bonds of a first subset as selected in (4) and atoms and/or bonds of a second subset as selected in (4), wherein the first subset and the second subset comprise atoms and/or bonds derived from opposite ends of the matching bond as selected in (4);
- (6) compare output ligands to the ligands of step (1);
- (7) store output ligands that are not identical to the ligands of step (1);
- (8) repeat steps (2)-(7), wherein the models superimposed in step (2) comprise the stored output ligands of step (7); wherein the repeating stops when each output ligand of step (7) is identical to a ligand provided in the previous step (2) of the repetition.

44. **(Previously presented)** An article comprising machine-readable media that stores executable instructions, the instructions causing a machine to:

- (1) map spatial relationships between two or more models of ligands of a set such that the models are superimposed, wherein each ligand comprises a plurality of atoms and a plurality of bonds, and wherein each model comprises three-dimensional structural information for a ligand; and
wherein the structural information is derived from physical observation and/or computational inference;
- (2) identify one or more pairs of matching bonds between ligands of the set, wherein the matching bonds comprise a bond of a first ligand (B1) and a bond of a second ligand (B2) that are superimposed in step (2) such that
 - (i) an atom at each end of the bond (B1) is within 1.8 angstrom of an atom at each end of the bond (B2),
 - (ii) the bond (B1) and the corresponding bond (B2) are of the same bond order, and
 - (iii) the bond (B1) and the corresponding bond (B2) are related by an angle of 30° or less;
- (3) select a plurality of subsets of atoms and/or bonds from each ligand;
wherein each subset comprises a bond and/or, an atom connected to a matching bond as identified in (2);
- (4) generate and display output ligands, each output ligand comprising atoms and/or bonds of a first subset as selected in (3) and atoms and/or bonds of a second subset as selected in (3), wherein the first subset and the second subset comprise atoms and/or bonds derived from opposite ends of the matching bond as selected in (3).

45. **(Previously presented)** The method of any one of claims 1, 39, and 42 to 44, wherein the structural information derived from physical observation is derived from X-ray crystallography or NMR.

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46. (Currently amended) The method of any one of claims 1, 39, and 42 to 44, wherein the structural information derived from computational inference is derived by modeling the structure of a ligand in a target protein macromolecule using computational means.